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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
10/661,966 09/11/2003		Richard B. Roth	SEQ-4038-UT	9006
47328	7590 05/03/2006		EXAMINER	
BIOTECHN C/O PORTFO	OLOGY LAW GROUP	SITTON, JEHANNE SOUAYA		
PO BOX 52050			ART UNIT	PAPER NUMBER
MINNEAPOI	LIS, MN 55402	1634	1634	

DATE MAILED: 05/03/2006

Please find below and/or attached an Office communication concerning this application or proceeding.

		Application	n No.	Applicant(s)				
Office Action Summary		10/661,96	6	ROTH ET AL.				
		Examiner		Art Unit				
		Jehanne S	S. Sitton	1634				
Period fo	The MAILING DATE of this communication or Reply	appears on the	cover sheet with the c	orrespondence a	ddress			
WHIC - Exter after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR RECHEVER IS LONGER, FROM THE MAILING asions of time may be available under the provisions of 37 CF SIX (6) MONTHS from the mailing date of this communication period for reply is specified above, the maximum statutory pere to reply within the set or extended period for reply will, by steeply received by the Office later than three months after the med patent term adjustment. See 37 CFR 1.704(b).	G DATE OF TH R 1.136(a). In no even n. eriod will apply and witatute, cause the appl	IIS COMMUNICATION ent, however, may a reply be timed Il expire SIX (6) MONTHS from ication to become ABANDONE	N. nely filed the mailing date of this of D (35 U.S.C. § 133).	•			
Status								
1)⊠	Responsive to communication(s) filed on 1	1 September 2	003.					
2a) <u></u>	This action is FINAL . 2b) ☐ This action is non-final.							
3)								
	closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.							
Dispositi	on of Claims							
4)⊠	4)⊠ Claim(s) <u>1-48</u> is/are pending in the application.							
	4a) Of the above claim(s) is/are withdrawn from consideration.							
5)	Claim(s) is/are allowed.							
6)□	Claim(s) is/are rejected.							
· · · —	7) Claim(s) is/are objected to.							
8)⊠	Claim(s) <u>1-48</u> are subject to restriction and	l/or election req	uirement.					
Applicati	on Papers							
9)	The specification is objected to by the Exan	niner.						
10) ☐ The drawing(s) filed on is/are: a) ☐ accepted or b) ☐ objected to by the Examiner.								
	Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).							
Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).								
11)	The oath or declaration is objected to by the	e Examiner. No	te the attached Office	Action or form P	TO-152.			
Priority u	ınder 35 U.S.C. § 119							
12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of:								
	1. Certified copies of the priority documents have been received.							
	2. Certified copies of the priority documents have been received in Application No							
	3. Copies of the certified copies of the priority documents have been received in this National Stage							
	application from the International Bu	•	` ''					
* 5	See the attached detailed Office action for a	list of the certi	fied copies not receive	ed.				
Attachmen	• •							
	e of References Cited (PTO-892) e of Draftsperson's Patent Drawing Review (PTO-948	1	4) Interview Summary Paper No(s)/Mail Da					
3) 🔲 Infor	e of Draitsperson's Patent Drawing Review (P10-946) nation Disclosure Statement(s) (PTO-1449 or PTO/SE r No(s)/Mail Date		5) Notice of Informal P 6) Other:		O-152)			

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DETAILED ACTION

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Election/Restrictions

- 1. Restriction to one of the following inventions is required under 35 U.S.C. 121:
 - I. Claims 1-15 drawn to a method for identifying a subject at risk of melanoma via the detection of polymorphic variation, classified in class 435, subclass 6.
 - II. Claims 16-21, drawn to a method for identifying a polymorphic variation associated with melanoma proximal to an incident polymorphism, classified in class 435, subclass 6.
 - III. Claims 22-25, drawn to an isolated nucleic acid comprising a polymorphic variation, classified in class 536, subclass 23.1.
 - IV. Claim 26, drawn to a polypeptide, classified in class 530, subclass 350.
 - V. Claims 27-33, drawn to a method for identifying a candidate test molecule that modulates cell proliferation, classified in class 514, subclass 1.
 - VI. Claims 34-36, drawn to a method of treating melanoma by contacting one or more cells with a nucleic acid, such as RNA, classified in class 536, subclass 24.5.
 - VII. Claim 37, drawn to a method of treating melanoma by contacting one or more cells with a protein, classified in class 514, subclass 2.
 - VIII. Claims 38-41, drawn to a method of treating melanoma comprising detecting a polymorphic variation, classified in class 514, subclass 1.
 - IX. Claims 42-44, drawn to a method of preventing melanoma by detecting a polymorphic variation and administering a melanoma preventative, classified in class 435, subclass 6.

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Claims 45-48, drawn to a method of targeting information, classified in class 702,
 subclass 20.

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2. Additionally, each group named above is subject to further restriction.

For Groups I, III, and IV, Applicant is required to further elect a specific SEQ ID NO (claim 1, 22) and to further elect a specific polymorphism or combination of polymorphisms to which the claims directed to specific polymorphisms will be limited. Additionally, for claim 14 or 24, applicant is required to elect a corresponding oligonucleotide.

For Group II, V, VIII, IX, and X, Applicant is required to further elect a specific SEQ ID NO (ie, claim 16, 27, 33, 38, 42, 45) and to further elect a specific polymorphism or combination of polymorphisms to which the claims directed to specific polymorphisms will be limited.

For Group VI and VII, Applicant is required to further elect a specific SEQ ID NO (ie, claim 34, 37)

This is NOT an election of species. The nucleotide sequences are structurally distinct chemical compounds and are unrelated to one another. These sequences are thus deemed to normally constitute independent and distinct inventions within the meaning of 35 U.S.C. 121.

Absent evidence to the contrary, each such nucleotide sequences are presumed to represent an independent and distinct invention, subject to restriction requirement pursuant to 35 USC 121 and 37 CFR 1.141. By statute, "[i]f two or more independent and distinct inventions are claimed in one application, the Commissioner may require the application to be restricted to one of the inventions." 35 U.S.C. 121. Pursuant to this statute, the rules provide that "[i]f two or more independent and distinct inventions are claimed in a single application, the examiner in his action shall require the applicant... to elect that invention to which his claim shall be restricted." 37

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CFR 1.142 (a). See also 37 CFR 1.141(a). It is noted that searching more than one of the claimed patentably distinct sequences and variants represents a serious burden for the office. The search for each sequence and variant are not coextensive and are required to be searched separately in both computer databases as well as nucleotide variant databases such as dbSNP.

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It is noted that any claim which specifically lists only a non elected sequence or variant will be withdrawn from consideration as being drawn to a non elected invention. Claims which are drawn to sequences and/or variants in the alternative and which include the elected sequence and/or variant, will be searched and examined with regard to the elected invention. If only one polymorphism is elected, claims directed to two or more polymorphisms will be withdrawn from consideration as being drawn to an non elected invention, but may be rejoined if they contain the limitations of the allowable single polymorphism. However, should the rejoinder require new grounds of rejection and occur after non final rejection, finality of the subsequent office action will not be affected. Amendments submitted after final rejection are governed by 37 CFR 1.116

It is noted that the claims refer to either a SEQ ID NO: or to a figure. Applicant is requested to indicate which SEQ ID NO is elected. If a SEQ ID NO: is not appropriate, ie: the figure refers to a snp or rs number, applicant is requested to indicate which variant is being elected, and if possible to indicate which SEQ ID NO: it is located in. Additionally, applicant should amend the claims to reflect the election.

3. The inventions are distinct, each from the other because of the following reasons:

The inventions of groups I-II, VIII, IX, and X are directed to related processes. The related inventions are distinct if the inventions as claimed do not overlap in scope, i.e., are

prevention, or information targeting, and vice versa.

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mutually exclusive; the inventions as claimed are not obvious variants; and the inventions as claimed are either not capable of use together or can have a materially different design, mode of operation, function, or effect. See MPEP § 806.05(j). In the instant case, the methods include steps of determining polymorphisms, however, the methods of group I of identifying a subject at risk of melanoma, vs group II, a method of identifying an incident polymorphism, vs group VIII, a method of treating melanoma, vs group IX, a method of preventing melanoma, vs group X, a method of targeting information, are not coextensive in scope as each method is directed to different steps to perform the method, ie: analysis of melanoma, analysis of another proximal polymorphism, treatment based on results of genotype analysis, prevention based on results of genotype analysis, and directing information. Further, the methods are not obvious variants. Each method has a materially different mode of design, mode of operation, function, and effect. A search for each of the patentably distinct process presents a serious search burden as the searches are not coextensive. Art directed to methods of identifying a risk of melanoma (group

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The inventions of groups I, II, and X are unrelated to the inventions of groups V - VI.

Inventions are unrelated if it can be shown that they are not disclosed as capable of use together and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06).

In the instant case, the different inventions have different designs, modes of operation and effects.

I), would not necessarily provide any information regarding proximal polymorphism, treatment,

The inventions of groups I, II, and VIII-X are unrelated to the invention of group IV.

Inventions are unrelated if it can be shown that they are not disclosed as capable of use together

and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06). In the instant case, the different products are not disclosed as capable of use with the methods and the different inventions have different designs, modes of operation and effects.

The inventions of groups I, II, and IX-X are unrelated to the invention of group VII.

Inventions are unrelated if it can be shown that they are not disclosed as capable of use together and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06).

In the instant case, the different products are not disclosed as capable of use with the methods and the different inventions have different designs, modes of operation and effects.

The inventions of group III and groups I, II, VI, VIII, IX, & X are related as product and process of use. The inventions can be shown to be distinct if either or both of the following can be shown: (1) the process for using the product as claimed can be practiced with another materially different product or (2) the product as claimed can be used in a materially different process of using that product. See MPEP § 806.05(h). In the instant case the products can be used to encode polypeptides or to detect expression, which are not required to practice the method of Group I or VII-X. Further, the product of group III is not required to practice the method of group II. The method of detecting a proximal polymorphism is not dependent on the detection of the incident polymorphism. The search for each group presents a serious search burden as the searches for each are not coextensive in scope. Art relating to polynucleotides would not necessarily provide any descriptive information regarding melanoma and vice versa.

The inventions of group IV and group VII are related as product and process of use. The inventions can be shown to be distinct if either or both of the following can be shown: (1) the process for using the product as claimed can be practiced with another materially different

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product or (2) the product as claimed can be used in a materially different process of using that product. See MPEP § 806.05(h). In the instant case the product can be used to make a fusion protein with enzymatic function, which is not required to practice the method of Group VII. Art relating to the structural requirements of the polypeptide would not necessarily have any information regarding treatment of melanoma and vice versa.

The inventions of groups III and IV are patentably distinct because they are drawn to different products having different structures and functions. The nucleic acid of group III is composed of deoxyribonucleotides linked by phosphodiester bonds and assumes the form of a double helix. The polypeptide of group IV is composed of amino acids linked by peptide bonds and can assume complex tertiary structures. The products of groups III and IV can be used in materially different processes, for example the DNA of group III can be used in hybridization assays and the polypeptide of group IV can be used to make a fusion protein with an enzymatic function. Consequently, the reagents, reaction conditions, and reaction parameters required to make and use each invention are different. Therefore, the inventions of groups III and IV are patentably distinct from each other. The search for each of groups III and IV presents a serious search burden as the searches for each are not coextensive in scope. The inventions have different status in the art as shown by their different classifications. In cases such as this one where descriptive sequence information is provided, the sequences are searched in appropriate databases. There is a search burden also in the non-patent literature. Prior to the concomitant isolation and expression of the sequence of interest there may be journal articles devoted solely to polypeptides which would not have described the polynucleotide. Similarly, there may have

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been "classical" genetics papers which had no knowledge of the polypeptide but spoke to the gene. Searching, therefore is not coextensive and presents a serious burden.

The invention of group III is unrelated to the invention of group V and VII. Inventions are unrelated if it can be shown that they are not disclosed as capable of use together and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06). In the instant case, the products are not used in the method of group V or VII and the different inventions have different designs, modes of operation and effects.

The invention of group IV is unrelated to the inventions of groups I, II, and V-VI, VIII-X. Inventions are unrelated if it can be shown that they are not disclosed as capable of use together and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06). In the instant case, the products is not used in the methods of groups I, II, and V-VI, VIII-X and the different inventions have different designs, modes of operation and effects.

The inventions of groups V-VIII are directed to related processes. The related inventions are distinct if the inventions as claimed do not overlap in scope, i.e., are mutually exclusive; the inventions as claimed are not obvious variants; and the inventions as claimed are either not capable of use together or can have a materially different design, mode of operation, function, or effect. See MPEP § 806.05(j). In the instant case, the methods include steps of treatment or modulation, however, the different methods: that is, group V, of identifying a modulator of cell proliferation, vs group VI, a method of treatment using nucleic acid, vs group VII, a method of treatment by administering a polypeptide, and group VIII, a method of treatment with surgery or chemotherapy, are not coextensive in scope as each method is directed to different steps requiring different reagents and parameters. Further, the methods are not obvious variants. Each

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method has a materially different mode of design, mode of operation and function. A search for each of the patentably distinct process presents a serious search burden as the searches are not coextensive. Art directed to each of the different modes of treatment or modulation would not necessarily provide any information on polymorphism or any of the other claimed modulators.

The inventions of groups V-VII are unrelated to the inventions of groups IX-X. Inventions are unrelated if it can be shown that they are not disclosed as capable of use together and they have different designs, modes of operation, and effects (MPEP § 802.01 and § 806.06). In the instant case, the different inventions have different designs and modes of operation and effects.

4. Claim 38 is generic to the following disclosed patentably distinct species in claim 41. The species are independent or distinct because they are drawn to structurally distinct therapeutics. Applicant is required under 35 U.S.C. 121 to elect a single disclosed species, even though this requirement is traversed. Applicant is advised that a reply to this requirement must include an identification of the species that is elected consonant with this requirement, and a listing of all claims readable thereon, including any claims subsequently added. An argument that a claim is allowable or that all claims are generic is considered nonresponsive unless accompanied by an election.

Upon the allowance of a generic claim, applicant will be entitled to consideration of claims to additional species which depend from or otherwise require all the limitations of an allowable generic claim as provided by 37 CFR 1.141. If claims are added after the election, applicant must indicate which are readable upon the elected species. MPEP § 809,02(a).

5. Because these inventions are independent or distinct for the reasons given above and have acquired a separate status in the art in view of their different classification, restriction for examination purposes as indicated is proper.

- 6. Because these inventions are independent or distinct for the reasons given above and the inventions require a different field of search (see MPEP § 808.02), restriction for examination purposes as indicated is proper.
- 7. Because these inventions are independent or distinct for the reasons given above and have acquired a separate status in the art because of their recognized divergent subject matter, restriction for examination purposes as indicated is proper.
- 8. Applicant is advised that the reply to this requirement to be complete must include (i) an election of a species or invention to be examined even though the requirement be traversed (37 CFR 1.143) and (ii) identification of the claims encompassing the elected invention.

The election of an invention or species may be made with or without traverse. To reserve a right to petition, the election must be made with traverse. If the reply does not distinctly and specifically point out supposed errors in the restriction requirement, the election shall be treated as an election without traverse.

Should applicant traverse on the ground that the inventions or species are not patentably distinct, applicant should submit evidence or identify such evidence now of record showing the inventions or species to be obvious variants or clearly admit on the record that this is the case. In either instance, if the examiner finds one of the inventions unpatentable over the prior art, the evidence or admission may be used in a rejection under 35 U.S.C.103(a) of the other invention.

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9. Applicant is reminded that upon the cancellation of claims to a non-elected invention, the inventorship must be amended in compliance with 37 CFR 1.48(b) if one or more of the currently named inventors is no longer an inventor of at least one claim remaining in the application. Any amendment of inventorship must be accompanied by a request under 37 CFR 1.48(b) and by the fee required under 37 CFR 1.17(i).

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10. The examiner has required restriction between product and process claims. Where applicant elects claims directed to the product, and a product claim is subsequently found allowable, withdrawn process claims that depend from or otherwise include all the limitations of the allowable product claim will be rejoined in accordance with the provisions of MPEP § 821.04. Process claims that depend from or otherwise include all the limitations of the patentable product will be entered as a matter of right if the amendment is presented prior to final rejection or allowance, whichever is earlier. Amendments submitted after final rejection are governed by 37 CFR 1.116; amendments submitted after allowance are governed by 37 CFR 1.312.

In the event of rejoinder, the requirement for restriction between the product claims and the rejoined process claims will be withdrawn, and the rejoined process claims will be fully examined for patentability in accordance with 37 CFR 1.104. Thus, to be allowable, the rejoined claims must meet all criteria for patentability including the requirements of 35 U.S.C. 101, 102, 103, and 112. Until an elected product claim is found allowable, an otherwise proper restriction requirement between product claims and process claims may be maintained. Withdrawn process claims that are not commensurate in scope with an allowed product claim will not be rejoined.

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See "Guidance on Treatment of Product and Process Claims in light of In re Ochiai, In re

Brouwer and 35 U.S.C. § 103(b)," 1184 O.G. 86 (March 26, 1996). Additionally, in order to

retain the right to rejoinder in accordance with the above policy, Applicant is advised that the

process claims should be amended during prosecution either to maintain dependency on the

product claims or to otherwise include the limitations of the product claims. Failure to do so

may result in a loss of the right to rejoinder.

11. Any inquiry concerning this communication or earlier communications from the

examiner should be directed to examiner Jehanne Sitton whose telephone number is (571) 272-

0752. The examiner can normally be reached Monday-Thursday from 8:00 AM to 5:00 PM and

on alternate Fridays.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's

supervisor, Ram Shukla, can be reached on (571) 272-0735. The fax phone number for this

Group is (571) 273-8300.

Any inquiry of a general nature or relating to the status of this application or proceeding

should be directed to (571) 272-0547.

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For all other customer support, please call the USPTO Call Center (UCC) at 800-786-9199.

Jehanne Sitton

Primary Examiner

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5/1/06